1 Claims

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- A method of producing an oligopeptide product,
- 4 the method comprising the steps:
- 5 a) providing a first oligopeptide, the first
- 6 oligopeptide having a reactive moiety,
- 7 b) providing a second oligopeptide, the second
- 8 oligopeptide having a activated ester moiety
- 9 c) allowing the reactive moiety of the first
- 10 oligopeptide to react with the activated ester
- 11 moiety of the second oligopeptide to form an
- 12 oligopeptide product, in which the first and second
- oligopeptides are linked via a linking moiety having
- 14 Formula I, Formula II or Formula III.

15

16 Formula I

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18 Formula II

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20 Formula III

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- 24 2. The method according to claim 1 wherein the
- 25 terminal activated ester moiety is a thioester
- 26 wherein the peptide is the acyl substituent of

1 the thioester.

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- 3 3. The method according to claim 2, wherein said
- 4 second polypeptide is generated by thiol reagent
- 5 dependent cleavage of a precursor molecule, said
- 6 precursor molecule comprising a second oligopeptide
- 7 fused N-terminally to an intein domain.

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- 9 4. A method of producing an oligopeptide product,
- 10 the method comprising the steps:
- 11 a) providing a first oligopeptide, the first
- 12 oligopeptide having a reactive moiety,
- (i) providing a precursor oligopeptide molecule, the
- 14 precursor oligopeptide molecule comprising a second
- oligopeptide fused N-terminally to an intein domain
- 16 (ii) allowing thiol reagent dependent cleavage of
- 17 the precursor molecule to generate a second
- 18 oligopeptide molecule, said second oligopeptide
- 19 molecule having a thioester moiety at its C-
- 20 terminus,
- 21 c) allowing the reactive moiety of the first
- 22 oligopeptide to react with the second oligopeptide
- 23 molecule to form an oligopeptide product, in which
- 24 the first and second oligopeptides are linked via a
- 25 linking moiety having Formula I, II or III.

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- 5. The method according to any one of the preceding
- 28 claims wherein the reactive moiety is a hydrazine
- 29 moiety, a hydrazide moiety or an aminooxy moiety.

- 31 6. The method according to claim 5, wherein the
- 32 reactive moiety is an aminooxy moiety and the

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1	activated ester moiety is a thioester.
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3	7. The method according to claim 5, wherein said
4	first oligopeptide is produced by reaction of
5	hydrazine with a precursor molecule, said
6	precursor molecule comprising a precursor
7	oligopeptide fused N-terminally to an intein
8	domain via a thioester moiety.
9	·
10	8. A method of producing an oligopeptide product,
11	said method comprising the steps:
12	a) providing a first oligopeptide, the first
13	oligopeptide having a reactive moiety, wherein
14	the reactive moiety is a hydrazine moiety, a
15	hydrazide moiety or an amino-oxy moiety;
16	(i) providing a precursor oligopeptide molecule,
17	the precursor oligopeptide molecule comprising a
18	second oligopeptide fused N-terminally to an
19	intein domain;
20	(c) allowing the reactive moiety of the first
21	oligopeptide to react with the precursor
22	oligopeptide molecule to form an oligopeptide
23	product, in which the first and second
24	oligopeptides are linked via a linking moiety
25	having Formula I, Formula II or Formula III.
26	
27	9. The method according to any one of the preceding
28	claims, wherein the first oligopeptide or the
29	second oligopeptide is a recombinant oligopeptide
30	and the other of the the first oligopeptide and
31	the second oligopeptide is a synthetic
32	polypentide.

1 2 The method according to any one of claims 1 to 3 8, wherein the first oligopeptide and the second oligopeptide are recombinant oligopeptides. 4 5 6 11. The method according to any one of claims 1 to 7 8, wherein the first oligopeptide and the second oligopeptide are synthetic oligopeptides. 8 9 A method of generating a protein hydrazide, 10 11 said method comprising the steps: 12 (a) providing a protein molecule comprising an 13 oligopeptide fused N-terminal to an intein 14 domain, 15 (b) reacting said protein molecule with 16 hydrazine, such that the intein domain is cleaved 17 from the oligopeptide to generate a protein 18 hydrazide. 19 20 The method according to any one of the claims 1 21 to 11 wherein step (c) of the method is performed 22 at a pH in the range pH 6.5 to 7.5. 23 A method of producing an oligopeptide product, 24 25 the method comprising the steps: 26 a) providing a first oligopeptide, the first 27 oligopeptide having an aldehyde or ketone moiety, 28 b) providing a precursor oligopeptide molecule, 29 the precursor oligopeptide molecule comprising a 30 second oligopeptide fused N-terminally to an

c) reacting said precursor oligopeptide molecule

intein domain,

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with hydrazine to generate an oligopeptide 1. 2 molecule comprising an intermediate oligopeptide, 3 said intermediate oligopeptide having a terminal 4 hydrazide moiety, 5 d) allowing the aldehyde or ketone moiety of the first oligopeptide to react with the hydrazide 6 7 moiety of the intermediate oligopeptide molecule to form an oligopeptide product, in which first 8 oligopeptide and the second oligopeptide are 9 linked via a hydrazone linking moiety. 10 11 An oligopeptide product produced by the method 12 of any one of the preceding claims. 13 14 15 A method of labelling an oligopeptide, the method comprising the steps: 16 a) providing a label molecule, the label molecule 17 having a reactive moiety, 18 b) providing the oligopeptide, the oligopeptide 19 having a activated ester moiety 20 21 c) allowing the reactive moiety of the label molecule to react with the activated ester moiety 22 of the oligopeptide to form the labelled 23 oligopeptide, in which the label molecule and the 24 oligopeptide are linked via a linking moiety 25 having Formula I, Formula II or Formula III. 26 27 17. The method according to claim 16, wherein in 28 step (c), where said label molecule and the 29 oligopeptide are linked via a linking moiety 30 having Formula II and where said activated ester 31 32 moiety of step (b) is not a thioester, said

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1	activated ester is a terminal activated ester
2	moiety.
3	
4	18. A method of labelling an oligopeptide, the
5	method comprising the steps:
6	a) providing a label molecule, the label molecule
7	having an activated ester moiety of which the
8	label is the acyl substituent,
9	b) providing the oligopeptide, the oligopeptide
10	having a reactive moiety
11	c) allowing the activated ester moiety of the
12	label molecule to react with the reactive moiety
13	of the oligopeptide to form the labelled
14	oligopeptide, in which the label molecule and the
15	oligopeptide are linked via a linking moiety
16	having Formula I, Formula II or Formula III,
17	wherein, in step (c), where said label molecule
18	and the oligopeptide are linked via a linking
19	moiety having Formula II and where said activated
20	ester moiety of step (b) is not a thioester, said
21	activated ester is a terminal activated ester
22	moiety.
23	
24	19. The method according to claim 18 wherein said
25	oligopeptide is produced by reaction of hydrazine
26	with a precursor molecule, said precursor
27	molecule comprising a precursor oligopeptide
28	fused N-terminally to an intein domain via a
29	thioester moiety.
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31 20. A method of labelling an oligopeptide, the 32 method comprising the steps:

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64 1 a) providing a label, the label having a reactive 2 moiety, 3 (i) providing a precursor oligopeptide molecule, the precursor oligopeptide molecule comprising an 4 5 oligopeptide fused N-terminally to an intein 6 domain 7 (ii) allowing thiol reagent dependent cleavage of 8 the precursor molecule to generate the oligopeptide molecule, said oligopeptide molecule 9 having a thioester moiety at its C-terminus, 10 11 c) allowing the reactive moiety of the label to 12 react with the oligopeptide molecule to form a 13 labelled oligopeptide, in which the label and oligopeptide are linked via a linking moiety 14 15 having Formula I, II or III. 16 The method according to any one of claims 16 to 17 21. 18 18, wherein the reactive moiety is an aminooxy 19 moiety and the activated ester moiety is a thioester. 20 21 22 22. The method according to claim 20, wherein the 23 reactive moiety is an aminooxy moiety. 24 A method of labelling an oligopeptide, the 25 23. 26 method comprising the steps: 27 a) providing a label molecule, the label molecule 28 having a reactive moiety, 29 b) providing a precursor oligopeptide molecule, 30 the precursor oligopeptide molecule comprising an 31 oligopeptide fused N-terminally to an intein

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domain,

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1	c) allowing the reactive moiety of the label
2	molecule to react with the precursor oligopeptide
3	molecule to form a labelled oligopeptide product,
4	in which the label molecule and the oligopeptide
5	are linked via a linking moiety having Formula I,
6	Formula II or Formula III as defined above.
7	
8	24. The method according to any one of claims 16 to
9	23 wherein step (c) of the method is performed at
10 <sup>.</sup>	a pH in the range pH 6.5 to pH 7.5.
11	
12	25. A method of labelling an oligopeptide, the
13	method comprising the steps:
14	a) providing a label molecule, the label molecule
15	having a aldehyde or ketone moiety,
16	b) providing a precursor oligopeptide molecule,
17	the precursor oligopeptide molecule comprising a
18	first oligopeptide fused N-terminally to an
19	intein domain,
20	c) reacting said precursor oligopeptide molecule
21	with hydrazine to generate an oligopeptide
22	molecule comprising an intermediate oligopeptide,
23	said intermediate oligopeptide having a terminal
24	hydrazide moiety,
25	d) allowing the aldehyde or ketone moiety of the
26	label molecule to react with the hydrazide moiety
27	of the intermediate oligopeptide molecule to form
28	a labelled oligopeptide product, in which the
29	label molecule and oligopeptide are linked via a
30	hydrazone linking moiety.
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L	26. The method according to claim 14 or claim 25,
2	wherein the aldehyde or ketone moiety is an $\alpha$ -
3	diketone or an α-keto-aldehyde group.
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5	27. A labelled oligopeptide produced by the method
5	of any one of claims 16 to 26.
7	